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(54) Title: COMPOSITIONS AND METHODS F TIVE AGENTS	OR TO	CAL ADMINISTRATION OF PHARMACEUTICALLY AC
(57) Abstract		

A composition for topical application comprising a therapeutically effective amount of a pharmaceutical agent(s), a flexible, finite, pharmaceutically acceptable, adhesive, and a solvent for the pharmaceutical agent(s) in the adhesive and a method of administrating the pharmaceutical agent to a mammal are diclosed.

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COMPOSITIONS AND METHODS FOR TOPICAL ADMINISTRATION OF PHARMACKUTICALLY ACTIVE AGENTS

CROSS-REFERENCE TO RELATED APPLICATION

This application is a continuation-in-part of U.S. Patent Application Serial Number 07/651,827 filed February 27, 1991, and U.S. Serial Number 07/813,196 filed December 23, 1991, both of which molications are hereby incorporated by reference.

Field of the Invention

The present invention relates compositions and methods for the administration of pharmaceutically active agents, namely those having a pharmacological or cosmetic effect, to a mammal in need thereof. The present invention is especially useful with local anesthetic agents for topical administration. In addition, the invention relates to a method for the topical administration of a pharmaceutical agent, especially an anesthetic agent or a combination of anesthetic agents, to prevent or ameliorate a disease or other medical or commetic condition, especially pain.

There is no limitation on the type of pharmaceutical agent that can be used in the present invention, provided that the agent can be absorbed percutaneously. Thus, the pharmaceutical agents can be drugs that can be topically applied for local effects and those which can be topically applied for systemic effects.

Background of the Invention

Anesthetic agents are pharmacologically active agents that block nerve conduction when applied in therapeutically effective amounts. They can be used for local or systemic effects. Anesthetic agents

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have been used extensively in the medical field to obtain topical anesthesia. Topical administration or application means the direct contact of the anesthetic with tissue to be anesthetized, such as skin or membrane, particularly the oral or boccal success. Previous methods of applying topical anesthetic sgents to the skin or mucosa have used "monfinite" or seniliquid carriers or spreading substances such as creams, eais or ointeents, or "finite" carriers, non-spreading substances which retain their form, e.g. patches, dressings and bandages. The finite carriers are flexible in the sense that they can bend to conform to the configuration of the skin or mucosa where they are applied.

Local anesthetics generally are esters or amides of bensoic acid derivatives, administered either as the free base or the acid-addition salt.

Free bases tend to be irritating at high concentrations. Acid-addition salts have low skin

permeability.

To be effective, a topical, local anesthetic should contain sufficient concentration of the active

smould comman suritisant consentation to a server agent to produce an anesthetic effect, it should penetrate intact skin or mucoss sufficiently to dailver a therapsutic dose, and it should exhibit rapid onset of anesthetic action and have a prolonged ansesthetic effect. In achieving the foregoing, it is often desirable to have the anesthetic agent present in a high concentration in the doseage form to effect a rapid onset and, additionally or elternatively, in excess of the amount that can be immediately absorbed through the duration or effect of anesthesia. On the other hand, the presence of the anesthetic agent in crystalline form may irritate sensitive tissues

such as mucosal tissues. This is particularly true

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vith regard to lidocaine. The usefulness of topical anesthetics has been limited by the concentration of drug achievable in the dosage form. The same considerations also apply generally to other bharmacoutically active acents.

Anesthetic agente have been used in monfinite form. United States Patent No. 4,894,232 to Redl, et al. discloses a base for mucosal or denture adhesive paates and a process for the preparation thereof. A lidocaine salt is named as suitable for this paste.

Finite local anesthetic compositions are reported in the literature. Some compositions are solvent free. For instance. Swedish Patent Publication No. 352,239 published December 27, 1972 in the name of S.G. Davis et al., assigned to Astra Pharmaceutical Products, Inc., and based on Swedish patent application No. 17744/70 filed December 30. 1970, disclosee a local anesthetic film containing up to 50% lidocaine in crystallized, microdispersed form. In its final form, this composition lacks a solvent for the anesthetic agent. The preparation is prepared by adding a solution of lidocaine in an organic solvent or an acid addition salt in water, under heat and agitation, to a solution or suspension of a filmforming material, namely carboxymethyl cellulose. polyvinyl alcohol, or a mixture of polyvinyl alcohol and polyvinyl pyrrolidone in water, followed by heating to remove any solvent present.

United States Patent No. 4,900,552 of Sanvoctaker et al., disclose a trilinainta fill suitable for prolonged and sustained delivery of an active ingredient in a buccal cavity. Specifically a hydratable succadeseive base layer, a non-adhesive reservoir layer containing the drug and a waterimpermeable carrier film sendviched between and bonded WO 92/15289 PCT/US92/01730

to the base layer and the reservoir layer form the trilaminate film.

Some finite anesthetic compositions contain polyhydric alcohol solvents. United States Patent Nos. 4,872,852 and 4,855,455 to Kigasawa and 3,249,109 to Maeth all describe the use of water soluble protein hased systems which incorporate anesthetics, and which also contain a tackifer and a polyhydric alcohol.

Some finite anesthatic agent compositions
on have a separate adhesive. United States Patent No.
3,814,095 to Lubens describes an absorbent pad for
topical application of an anesthatic agent having a
peripheral adhesive.

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olycerol (glycerin) has been used as a planticister for karvay gus. United States Patant Nos. 4,307,717 and 4,675,009 to Byses at. al., describe a druy in a solid phase formed of a synthatic polysacoharide or a combination thereof and a liquid phase of vater or an alcohol or a combination thereof. The amount of druy in the preparation (excluding solvent or carrier) is low. The cross-linked polysacoharide plasticised with water and/or a polyhydric alcohol is said to be not self-edmetring.

polypsaconaride plasticises with water analyse a polyhydric alcohol is said to be not self-adhering. The formulations do not include both a solvent for the drug and a plasticiser for the polysacoharide. It is also known to combine two local

anesthetic free bases with different melting points.

By mixing the two anesthetic bases, an entectic
mixture has been reported that is liquid at room
temperature, making it possible to attain higher
concentrations of the active bases. United States
Patent No. 4,885,554 to thang relates to a combination
of the free base and an acid addition salt or a
variety of drugs, typically in a liquid carrier, to
increase skin penetration rates. Amesthetics, along

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with a list of other suitable drugs are mentioned. This reference specifically teaches that base and acid-addition forms of the same drug be used in carrier.

United States Patent No. 2,352,691 to Curtis
teaches the use of salicylate salts of alkamine esters
of aminohenoic sold to enhance the water solubility
of ansethetic spents. In one example, this reference
discloses a solution of procaine acetyl salicylate
containing insoluble ansethetics such as bensocaine,
butesin, orthoform, or their salts, in certain
giycole, which are combined with a volatile solvant,
and then used to saturate gause bandages or other
mutable fabrics.

United States Patent No. 2,143,537 to Tieza describes an ointensat containing isosanyly/drocupyersies in combination with a quick acting local anesthetic to overcome the undestrable irritation caused by the prolonged acting anesthetic isosanyly/drocupyershe or its salts. The preparation of Tieza combines short and long acting anesthetic agents.

United States Patent No. 2,277,035 to Curtis relates to preparations containing a mixture of two or more anesthetic agent saits having different pH values in solution, whereby the pH value of the combined mixture in solution may be adjusted to obtain a higher degree of stability of the solution, and at relatively higher pH, a more rapid onset of anesthetic action. The anesthetic agents in Curtis are not in highly dispersed form and are used in a liquid-soaked fabric.

topical ensethetics has been schieved by the addition of vasoconstrictors, such as the catecholazine, epinephrine, which caused constriction of blood vassals. Since catecolazines are not particularly effective when applied topically, such a prolongation WO 92/15289 PCT/US92/01730

is of minimal usefulness for topical anesthetics. The primary drawbacks of this approach are the potential adverse side effects of categolamines, and the

prolongation itself.

Although many local ensethetic compositions have been proposed, it has been discovered that the incorporation of one or nore anesthetic agents in a solvent for the anesthetic agent or agents into a flexible, finite, pharmaceutically acceptable carrier, permits an exceptionally high locating of anesthetic agent in the carrier, permitting more rapid delivery of the anesthetic agent to the dermal suchemen and a greater extent of anesthesis without crystallization of the anesthetic agent for agents which can limit absorption by the skin and which can cause irritation of the asket or other dermal membrane.

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It has also surprisingly been found that concentrations of substantially dissolved anesthetic agent as high as 50° by weight of the total composition can be achieved in a system in which the adhesion of the adhesive is not hindered. Prolongation of anesthesia can thus be achieved by increasing the amount of time the composition is applied, without detrimental irritation.

The compositions of the present invention are in convenient form for topical application of the anesthatic agents, thereby enabling such anesthatics to penetrate the dermis, for example, intact skin or a succous membrane. Moreover, the anesthatic action is highly localized. Because the drug is substantially microdispersed in the carrier, it is more readily available for permestion into the skin or dermal

It still further has surprisingly been found that the use of two different local anesthetic agents, the first in base form and the second in acid-addition

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membrane.

salt form, in a finite, flexible, adhesive, pharascentically acceptable carrier, including a solvent for the anesthetic agents, paraits the attainment of anesthetic agent concentrations in the final product of up to 50¢ by weight in microdiapersed form, without crystallization of the anesthetic agents which can cause irritation of the ekin or other dermal

Thus, in one embodiment, the present invention is in convenient form for topical application of the amethetic agents, thereby emabling such anesthetics to penetrate intact skin or mucous membranes and have a highly localized effect. Furtherwore, the combination of the sait and base forms, advantageously results in rapid onset of amesthetic action with prolonged amenthatic effect.

Summary of the Invention

The invention relates to a flexible, finite

bioadhesive composition, for topical application
comprising:

a therapeutically effective amount of at least one local anesthetic or other pharmaceutically active agent which is in solid form at ambient temperatures and pressures;

a pharmaceutically acceptable solvent for the anesthetic or other pharmaceutically active agent, in an amount from about 5 to about 70 weight percent based on the weight of the whole composition, said solvent including about 5 to about 50 weight percent based on the weight of the whole composition of a planticiars for the bloadbastwe;

other pharmaceutically active agant in the solvent, a flexible, finite, pharmaceutically acceptable polysaccharide bioadhesive in an amount from about 20

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in admixture with the anesthetic agent or

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to about 50 weight percent based on the weight of the whole composition;

wherein the composition is substantially free of water, substantially water insoluble and selfadhesive; and wherein the pharmaceutically active agent is present in non-crystallized form in the

composition.

In another embodiment, the flexible, finite composition of the invention is comprised of two

anesthetic agents, that is:

a therapeutically effective amount of a first local anesthetic agent in base form;

a therapeutically effective amount of a different, second local anesthetic agent in acidaddition salt form;

a solvent for the first and second local amesthetic agents, preferably in an amount from about 5 to about 70 weight percent based on the weight of the whole composition; and

in an admixture with the anesthetic spants and the solvent, a pharamoeutically acceptable adhesive, preferably a bloadhesive, preferably in an amount from about 20 to about 50 weight percent based on the weight of the whole composition;

wherein the composition is preferably substantially free of vator, substantially water insoluble and selfadhesive; and wherein the anesthetic agents preferably are in non-crystallised form in the composition. The compositions of the invention may be

further include a backing material which conforms to the size and shape of a single dosage of the composition.

The present invention further relates to a method of administering one or more pharmaceutically active agents in a bloadhesive to a subject comprising the steps of:

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providing a composition comprising a therapeutically effective amount of at least one pharmaceutically active agent which is in solid form ambient temperatures and pressures: pharmaceutically acceptable solvent for pharmaceutically active agent, preferably in an amount from about 5 to about 70 weight percent based on the weight of the whole composition, said solvent preferably including about 5 to about 50 weight percent of a plasticizer for the bioadhesive; and in admixture with the pharmaceutically active agent in solvent, a pharmaceutically acceptable polysaccharide bioadhesive, preferably in an amount from about 20 to about 50 weight percent based on the weight of the whole composition; wherein said composition is substantially free of water, is substantially water insoluble and is self-adhesive; and wherein the pharmaceutically active agent is in non-crystallised form in the composition; and

contacting an area of skin or mucous membrane with the composition to administer the pharmaceutically active agent.

The invention further relates to a method of administering two local anesthetic agents to a subject comprising the steps of:

providing a composition comprising a therapsutically effective amount of a first local anesthatic agent in base form; a therapsutically effective amount of a different, second local amentatic agent in acid-addition salt form; a pharmaceutically acceptable solvent for the amentatic preferably in an amount which ranges from about 50 to about 70 weight percent based on the weight of the whole composition, said solvent preferably including about 5 to about 50 weight percent of a planticiser for the bloodheavie carrier; and in additure with the

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pharmaceutically active agant in the solvent, a pharmaceutically acceptable preferably polysaccharide bloadhesive, preferably in an amount from about 20 to about 50 weight percent beased on the weight of the whole composition; wherein said composition is preferably substantially free of water, substantially water inscludes and self-adhesive; and wherein the pharmaceutically active spent is in non-crystallised form in the composition; and

contacting an area of skin or mucous membrane with the composition thereby administering the local anesthetic agent.

The compositions of this invention permit a far higher leading of drug than conventional desage forms. This leading in the case of anesthetic agents can result in an extent (depth) of anesthesia which numbs the teeth when applied buccally, not a typical result for a topical anesthetic cream or ointeent.

Detailed Description of the Invention

This invention provides a composition which adheres to an area of the skin or mucosa, and permits delivery at elevated levels of pharmaceutical agent or a combination-of agents to produce a local or systemic effect over a prolonged period of time.

In accordance with one embodisent of the present invention, a local anesthetic in solution with a solvent for the anesthetic, containing a plasticizer for the admesive, is in admixture with a pharamecutically acceptable admesive, which is preferably a bicadhesive, and more preferably a polysacchariche bicadhesive, is provided in a finite, flexible form for topical application to the skin or dermal membrane of a mamma.

In accordance with a further embodiment of the present invention, a combination of local anesthetic agents, a solvent for the anesthetic agents

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and a flexible, preferably adhesive pharmaceutically acceptable adhesive carrier is provided for topical application to the skin or mucosa of a mammal.

The anathatic agents of this invention are those known, or of a type known, in the art. The local anesthetic bases encompassed by this invention are weak organic bases which are lipophilic in nature and thus poorly soluble in water. However, these bases will react with organic or inorganic acids to form acidic, water soluble acid-addition salts.

The base form and the sait form of the anesthetic agent incorporated in the combination composition of this invention must be different anesthetic agents, to achieve maximum duration of the anesthatic effect. By the term "different" is meant that that all form in any combination is not a sait of the base form used in the given combination.

Local anesthetic spents suitable for use in the practice of this invention include antides and estars. Examples of tha anides are lidocalms, princeains, sepivacains, buyinvacins, dibucains and atidocains. Esters include procains, tetracains, propoxycains, chloroprocains, benacoains, butamben picrats, cocalins, baylcains, piperocains, oxyprocains and proparacsine. Other suitable local anesthatics for use in the practice of this invention include cyclomethyanis, dimethisequin, ketocains, disperdon, dyclonine and praesoxins, all typically administered in the form of the acid addition hydro-chloride or suitates saits.

The acid-addition salts of the present invention are any non-toxio, pharmaceutically acceptable organic or inorganic salts. Typical inorganic salts are the hydrogen halides, especially the hydrochlorides, carbonates, borates, phosphates, sulfates, hydropromises, nitrates,

sulfides, and arsenates. Typical organic salts are salts of mono- and polycarboxylio acids such as the citrate, tartrate, malate, cinnamate, oxalate, formate, succinate and phthalates.

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The solvents for the anesthetic agents or other drugs are non-toxic, pharmaceutically acceptable substances, preferably liquids, which do not adhesion substantially negatively affect the properties of the system and in which the anesthetic agents or other drugs in the amounts employed are fully soluble. Preferably, the solvent is or is primarily a polyhydric alcohol or combination of polyhydric alcohols, particularly when the adhesive is a gum. The term polyhydrio alcohol means any organic polyol. Other suitable solvents include carboxlyio acids and their derivatives and analogs such as fatty acids such as cleic acid, linoleic acid, capric acid and the like, as well as fatty esters or alcohols and ketones such as polyvinylpyrrolidone.

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ketones such as polyvinylypyrolidones. Furranz suitable solvents include other non-toxic, nonvolatile solvents commonly used in decreal or transdurmal compositions for dissolving like compounds. As apparent to one skilled in the art what is a suitable solvent varies with the solubility of the drum in question.

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The above mentioned polyhydrio alcohols may include those having 2 to 6 alcoholic hydroxyl groups. Such polyhydrio alcohols include glycols, tricls and polyols having 4 to 6 alcoholic hydroxyl groups. Typical of said glycols are glycols containing 2 to 6 carbon atoms, e.g. ethylans glycol, propylans glycol, butylens glycol, polyethylans glycol, average molecular weight about 200 - 8,000, preferably about 200 to 6,000), dipropylans glycol, havylans glycol, polyovyethylans, polypropylans glycol, berylans glycol, betwien gl

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trimethylolpropane. Said polyols are exemplified by cycloalkamepolyols such as polyols derived from monosaccharides such as scortict (scort)t. These polyhydric alcohols may be used either singly or in combination (preferably, of two or three). Thus, for example, glycerin alone or a mixture of glycerin and butylame glycol is employed. In general, when an anesthatic agent, especially an anesthetic base is used, there are limits to the amounts of lipophilic polyhydric alcohols containing some than two elocholic hydroxyl groups that can be present in the solvent and yet not result in precipitation of the drug as

crystals. Among those polyhydric alcohols, those which satisfy the requirements relevant to the adjustment and maintenance of softness of the external drug of the invention, the compatibility or co-dispersibility with the other components, and provide a proper consistency of the composition, may be freely used. Those which are low in volatility and plastic, are generally preferred and, in this regard, dipropylene glycol, glycerin, propylene glycol, butylene glycol, and sorbitol are appropriate solvents, according to the invention. Since solvent is to remain, at least in part, in the composition, the solvent should include components that do not substantially volatilize under the drying conditions used in preparing the composition. In other words, the solvent for the drug should be non-volatile.

Solvent selection for a single anesthetic agent or a combination of anesthetic agents in either the free base form or in the acid-addition salt form, depends on the form of the anesthetic agent, namely whether it is in free base form or coid-addition salt form. Solvents for the salt form of anesthetic agent are polar organic solvents. Polar organic solvents

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are preferably polyhydric alcohols, as discussed above. Various other solvents suitable for either these or sold-sedition form of the anesthetic agent are those solvents known to dissolve either or both or these two types of forms including cyclic ketomes such as 2-pyrrolidone, 18-(2-hydroxysthyl) pyrrolidone, Reschylpyrrolidone, 18-deceylasopyloheptan-1-one and other n-substituted alkyl-assopyloheptal-2-ones (asones) dischylforzadde, and disetylpulfoxide.

Other suitable solvents for the free base form of the aneshetic agent are cell envelope disordering compounds known to be useful in topical pharmaceutical preparation, which compounds are thought to assist in skin penetration by disordering the lipid structure of the stratum cornaum cell-envelopes. Some of these compounds are generally encompassed by the formula

P-X

Although the exact amount of the polyhydric alcohol or alcohole in the composition depends on the nature of other components, and therefore cannot be stated in specific terms, the proportion may range

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from about 5 to about 70 weight percent based on the whole composition.

The solvent includes from about 5% to about 50% and more preferably about 10% to about 30% of a polyhydric alcohol known to plasticize the bicachesive carrier. A particularly useful plasticizer is giverine.

The high concentrations of microdispersed drug, for example anesthetic egent, of this invention are achieved typically by mixing the amenthatic agents with the solvent, preferably at an elevated temperature, for example about 70 to 100°C, to obtain a mixture, preferably a solution, of the amenthatic agents which is then added to the pharmaceutically acceptable addesive.

Preferably the anesthetic sgent is substantially dissolved in the solvent so that when mixed with the adhesive, the anesthetic is microdispersed in the composition. The term "microdispersed" is intended to mean that in the solvent, and subsequently in the carrier, there is an initiated dispersion of the anesthetic agent at the molecular or, ionic level, such that crystals of the anesthetic agent cannot be detected using a microscope having a magnification of roughly 25%. As such, the pharasceutically active agent is in "mon-crystalized" form when in the compositions of the present invention.

It has been discovered that high concentrations of a combination of nicrodispersed anesthetic agents, namely up to 50% by weight of the finite, flexible composition, require the use of a solvent as harein described. Onlession of the solvent in the procedure of Example 1 below yields a product filled with crystals or crystalline mass.

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In particularly preferred embodiments of this invention, the free base local anesthetic agent is selected from the group comprising lidocalne, processine, proposycaine, meptweatne, prilocalne, dyclonine, premaxine, bennecaine and chloroprocalne. The salt form is preferably one selected from the group comprising prilocalne, tetracaine, buptweatne, dyclonine, dibucaine, etitocaine and lidocaine salts. The aforementioned bases and salts can be used alone or in combination with other memethetic bases and salts as needed to achieve therepeutically affective lavels when administered transdermally.

The term "therapeutically effective amount" is intended to mean the amount of drug as a minimizer sufficient to produce a therapeutic effect, for example, an anesthetic effect when applied topically. These amounts are known in the art or may be determined by methods known in the art, and typically range from about 1 to 20,000 mg per human adult and preferably about 10 to 10,000 mg and most preferably range from about 20 to 5,000 mg of the anesthetic agent per application, depending upon the anesthetic agents chosen, and whether the skin or nucous membrane is the site of action. The only upper limit on the amount of anesthetic in the composition is that the preparation is substantially free of crystals of anesthetic agent or other drug and the amount of solvent used is not sufficient to undesirably affect the adhesive properties of the whole composition. Thus, the single ingredient anesthetic agent contains as a minimizer a therapeutically effective amount of anesthetic agent within the foregoing range.

The concentration as well as the quantity of amesthetic per squars centimeter can be varied independently in order to achieve the desired effect. Higher concentrations of amesthetic base contained in

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a dosage form of decreased thickness will result in a anesthetic with fast onset and short duration. High concentrations of the anesthetic base contained in a dosage form of increased thickness (higher mg of anesthetic per square centimeter) will result in potent anesthesia with fast onset and long duration. Low concentrations of the anesthetic base in a dosage form of decreased thickness will result in mild anesthesia with longer onset and short duration. Low concentrations of the anesthetic base contained in a dosage form of increased thickness will have mild anesthesia with longer onset and longer duration. As shown in the above explanation, the ability to vary the concentration of anesthetic from very low (about 1%) to high (40% or higher) of the total composition, when combined with the ability to coat thin (about 0.001 inches) or thick (about 0.500 or more inches) enables the practitioner of the invention to vary the dosage of the system as needed for particular anatomical sites of interest.

As a general rule, in the case of mucosal application, the ansethatic drug selected, the concentration and thickness and the duration of the application is determined based upon the anesthetic's ability to penetrate the mucosa and to be at peak effectiveness within about 2 to 30 minutes. The duration of the effect of the anesthetic on the oral mucosa should range between about 2 to 240 minutes, depending on the anesthatic agent selected, the concentration of the anesthatic agent selected, the concentration of the anesthatic agent selected, the concentration to the anesthatic agent selected, the concentration confidence of the should be selected dependent on meed, as will be apparent to one skilled in the art.

The ratio of the free base form to the salt form in the alternate composition of this invention will depend on several factors, namely: (1) the

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identity of the sait and hase used; (2) the desired duration of action; and (3) the desired rapidity of anasthetic effect. As a general rule in the case of mucosal application, the ratios of hase to sait are such that the free hase form preferrably should penetrate the success and he at its peak effectiveness within about a 2 to 10 minute period, whereas, the sait form should preferably penetrate the success and he at its peak effectiveness within a period of about 10 to 75 minutes. The duration of the effect of these on the oral success will maps between about 2 to 240 minutes depending on the hase/sait combination selected and the length of soulication time.

The term "onset of ansethesis" is intended to mean the time to peak effect on the individual narves. Onset of ansethesia principally depends upon the lipid solubility, molecular size, and quantity of available, un-ionized form of the local anesthetic. Thus, ansethetics with a high lipid solubility or a low pK, or both, have a more rapid onset of engethesis.

The term "duration of anesthesia" as used herein means the period of time during which the local anesthetic measurably blocks nerve conduction. The foregoing depends upon all of the factors listed for onset of anesthesia, as well as on the extent of wrotein binding of the amenthesic agent.

The unsethetic agent free base can penetrate intact skin to a limited degree, and will more rapidly penetrate the skin if the keratin layers are abraded. In the case of the oral mucosa, the anesthetic base will penetrate much more readily due to the different keratin composition and the resulting difference in the hydrophilicity as compared to the giratum corneum of intact skin.

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As a general rule, the salt forms of the aforementioned anesthetics do not appreciably penetrate intact skin, but the un-ionized base form do penetrate to a limited degree. Both forms, salt and base, will penetrate abraded keratin layers. The salt as well as the base will penetrate, to a differing degree, the buccal mucosa due to the buccal mucosa's hydrophilicity, as compared to the stratum corneum of intact skin. Generally, the higher the lipid content of the mucosal membrane, the more rapidly the base form of the anesthetic agent will be absorbed. Therefore, when the composition is used for application to oral or buccal mucosa, the different lipid contents of the gum (gingiva) and the alveolar mucosa must be kept in mind in order to obtain the optimal penetration rate.

Although applicants do not intend to be bound by any theory or proposed senchanism of operation, it is believed that the base which is lipid soluble has a rapid onset of anesthesis aince it enters the lipo-protein nerve membrane preventing the depolarization and ion exchange involved in stimulus conduction. On the other hand, the salt which is not lipid soluble, penetrates to the lipo-protein nerve membrane only after the buffering capacity of the akin or mucosal tissue converts the salt to the base, the final result before a delayed onset of anesthesis.

The saits of this invention in the combination composition are exacted on the basis of onset of anesthesia and duration of anesthesia. Adjusting the ratio of base to sait affects the relative onset as well as the duration of anesthetic action. The greater the amount of anesthetic agent having a rapid onset of action, the shorter the onset of anesthesia. Similarly, the greater the amount of the anesthetic agent having a prolonged duration of

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anesthesis, the more prolonged the duration of anesthesis. Mora than two anesthetic agents may be used to have a broader spectrum of activity. Moraover, the composition can include other drugs used concemitantly.

Generally, the concentration of solubilised anesthetic agent can range, on a weight basis, between about 1 and about 50% or more, preferably between 2.5 and 40% and more preferably between 5 and 30% of the total weight of the composition. In a preferred embodiment of tha combination of this invention, the concentration of dissolved basa is 20% by weight of the total composition. The base used in the preferred embodiment for a single ingredient preparation is 140ccating.

cenerally, for the hydrochloride salts the ratio by weight of base to salt is about 90:10 to about 60:40, preferably about 79:25 to about 60:40, and more preferably about 70:30 to about 60:40. For other salts, the ratios are comparable based on relative solar amounts. In a preferred embodiment of the invention, the ratio is about 2:11 base to salt, respectively. The base used in the preferred embodiment is il lidocaine and the preferred salt is a salt of prilocaine, buptweenine, dyclomine, mapitwacaine, or tetracaine, preferably the hydrochloride salt.

Table 1 below summarizes the peak and duration of action of selected local anasthatics based primarily on application to skin or mucous membranes:

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TARLE 1

5	Local Anesthetic	Minimum Adult Dose	Maximum Adult Dose (mg)	Peak Effect (minutes)	Duration of Effect (minutes)
10	Dibucaine Lidocaine Benzocaine		25 750 5000	< 15 2-5 1	120-240 30-60 30-60
	Cocaine		50	2-5	30-120
	Tetracaine Dyclonine		50 100	3-8 < 10	30-60 < 60
15	Pramoxine		200	3-5	NA
	NA:	Not Avail	able.		
	Sources	Drug Facts	and Compan	rianna 10	on edition

J.B. Lippincott Company, St. Louis, MO.
Page 601.

In general, the relative speed of onset of ansethesia and duration of anesthesia for any given form of anesthetic agent is available in the literature or can be calculated by standard tests. Onset time, as well as duration of

anesthesia, will wary from individual to individual as well as on the basis of the site of application. When applying the composition to highly keratinized dermal tisusus, the onset of anesthesia may take as long as 2 to 4 hours.

The composition of this invention can be manufactured by numerous methods known in the art which permit the achievement of a microdispersed anseshetle agent, including attruding, molding, solvent casting, coating, and all other methods which employ a solvent to disperse the drug in a carrier prior to shaping of the carrier.

Contrary to the typical method for manufacturing a drug in a solvent containing adhesive, the preparation is either not dried so as to force removal of the solvent from the adhesive or a solvent

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is used which is not substantially evaporated during the conditions of manufacture. The composition in question can then be applied to a flexible backing or a combination of backings which will serve to define the size and shape of a single dosage of the composition. Such backing may be a three dimensional material such as paper, a non-worn fabric or natural or synthatic polymer substance. Methods of coating backings are well-known in the art and include techniques involving Mayer rod, gravure, and knife-over roll. Further processing of backings may involve the use of converting equipment for die cutting.

The finished dosage form will be substantially occlusive to water permeation in invivo.

For example, the anesthetic agents are dissolved in a solvent, preferably a polyhydric alcohol, and then the resulting mixture is added to an adhesive prior to being placed onto the flatible form or hacking. The final form in which the composition of the invention will be applied depends upon the anatomical site of supplication.

The phrase "flexible, finite" with reference to the pharmacoutically acceptable carrier, is intended to sean a solid capable of conforming to a surface with which it comes into contact and capable of maintaining the contact to as to facilitate topical application without any adverse physiological response, and which can be used to establish the compositions herein in their preferred solid form without being appreciably descomposed by aqueous contact during administration to a patient.

An important characteristic of the present invention relates to the substantially water-free and water-insoluble nature of the composition. By the term "substantially water-free" is meant that the

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preparation contains less than about 10% by weight water, and preferably less than 5%, and most preferably less than 3%. In general, it is desirable to avoid the addition of water entirely and to eliminate, as far as possible, the presence of water in the other ingredients of the composition. By the term "substantially water insoluble" is meant that the composition remains "finite" and does not generally detach from the skin or other dermal membrane at the site of application and under the conditions of regular, intended use for a period of at least 3 The advantages to be derived from the substantially water-free and water-insoluble nature of the compositions of the present invention include achievement of higher concentrations of drug. Another advantage of these compositions is minimization of precipitation of drug into crystals. precipitation affects processing of the composition, affects rate of delivery of the drugs and in certain cases can affect sensitivity of the subject to be treated to the drug.

Suitable adhesive carriers include any of the non-toxic polymers, particularly those of the type used to carry drugs for transdermal delivery including natural or synthetic elastomers, auch polvisobutylene, styrene, butadiene, styrene isoprene block copolymers, acrylics, urethanes, silicones, styrene butadiene copolymers, methyl acrylate copolymers, acrylic acid, polyacrylates, polysacchrides such as, karaya gum, tragacanth gum, quar qum, cellulose, and cellulose derivatives such as methyl cellulose. cellulose, cellulose acetate and the like, along with other substances known for use in transdermal preparations capable of forming a solid colloid that can adhere to skin and mucosa, used alone or in

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combination with other suitable carriers. A particularly preferred carrier is a bloadhesive and more preferably a polysaccharide bloadhesive for application to the dermis, preferably the success. 2 adhesive on he modified so as to adhere to the skin or success tissue, depending on the intended amplication site.

The term "adhesive" as used herein means a substance, inorganic or organic, natural or synthetic, that is capable of surface attachment to the intended application site.

The term "bloodhesive" as used herein means an adhesive which attaches and preservably strongly attaches to a live or freshly killed biological surface such as skin or success! tissue upon hydration. Indeed, to qualify as a bloodhesive, a substance sust be capable of maintaining adhesion in noist or set in in-vive or in-vitro environments. The final composition of the present invention is "self-adhesive" in that it attaches to the site of interest vithout the need to reinforce its attachment by way of another adhesive which is applied to the composition.

The strength of adherence can be measured by standard tests for measuring the force, e.g., in dynes per square centineter, as disclosed in U.S. 4,615,697. Suitable bloadhesives include those prepared from optionally partially esterified or etherified to, polyacrylic acid polyaers, including but not limited to, polyacrylic acid polyaers lightly cross-linked with a polyalkenyl polyether or other ross-linking agent such as those commercially available from S.F. Goodrich, cincinanti, ohlo, under the trademarks Carbopol 394, 9349, 940 and 941.

Other suitable bioadhesives include natural or synthetic polysaccharides. The term "polysaccharide" as used herein means a carbohydrate

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decomposable by hydrolysis into two or more molecules of natural or synthetic moneascoharides or their analogs or derivatives. Suitable polysaccharides include cellulose derivatives such as methylcelullose, cellulose acetate, carboxymethylcelullose, hydrolysethylcelullose and the like. Other suitable bloadhesives are pactin, a sixture of suitable bloadhesives are pactin, a sixture of suitable such as natural plant exudates, including karaya gum, phatti gum, tragacanth gum, sundang mum, jaraya gum and the like, as well as esed gums such as guar gum, locust bean gum, postati plant eas guar gum, locust bean gum partini gum and the like.

In addition to the above ingredients, there asy also be incorporated other additives eslected from among the various pharmaceutically acceptable additives available to those skilled in the art. These additives include binders, stabilizers, preservatives, penetration enhancers, flavorings and pigmants. In the preferred embodiment, the compositions of the present invention also contain a binder or emulsifier such as lecithin which promotes dispersion of the other ingredients having differing solubilities, thereby enhancing the uniform consistency of the final commodition.

The composition is administered in appropriate since, typically having a surface area of from about 0.1 to about 200 cm² or conveniently 0.2 to 100 cm². The anesthetic agent is loaded into the composition in as high a concentration as necessary to effect therapy, e.g., in a range from about 0.1 mg/cm² to about 50 or more mg/cm².

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		position can	have the
following types and an	nounts of	ingredients:	
Ingredient	Typical Range (% by weight)	Preferred Range (% by weight)	Optimum Range (% by weight)
Adhesive	15 to 60	20 to 50	20 to 35
solvent (plasticizer included in solvent)	2 to 75 1 to 50	5 to 70 5 to 50	20 to 40 10 to 30
Anesthetic agent (single ingredient)	1 to 50	5 to 40	10 to 30
Anesthetic agent (multiple ingredient)	1 to 50	5 to 40	10 to 30
(a) Anesthetic base (b) Anesthetic salt	.7 to 50 .3 to 25	5 to 40 2 to 30	7 to 20 3 to 20
		the flexible	
bloadhesive composit	ion for	topical ap	plication
comprises:			
a therapeut	ically ef	fective amou	ant of at
least one pharmaceutic			
solid form at ambient			
		cceptable so	
the pharmaceutically a			
the pharmaceutically a about 5 to about 70 wei			
about 5 to about 70 well of the whole composition	dur barce	almont include	ling about
of the whole composition	on, sala s	Olvelic Illoin	m for the
5 to about 50 weight p	ercent or	a plasticize	it tor che
bioadhesive;			
		the pharmac	
active agent in the	solvent	, a pharmac	
acceptable polysaccha	ride bioa	dhesive in	an amount
from about 20 to about	50 weigh	t percent bas	ed on the
weight of the whole co	mposition	1	
wherein the composit:	ion is s	ubstantially	free of
water, substantiall	y water	insoluble a	nd self-
adhesive; and wherein	the ph	armaceutical	y active
meetic, and anoron			

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agent is present in non-crystallized form in the composition.

In another embodiment, the flexible, finite composition of the invention comprises:

a composition for topical application comprising a therapsutically effective amount of a first local anesthetic agent in base form and a therapsutically effective amount of a different, second local anesthetic agent in salt form in a pharmacoutically acceptable, adhesive-containing carrier containing asolvent for the first and second local anesthetic agents.

wherein the composition is preferably substantially free of water, and substantially water insoluble and is self-adhesive; and wherein the anesthetic agents are in non-crystallized form in the composition.

Preferably, the pharmaceutically acceptable solvent is in an amount from about 20 to about 53 weight percent based on the weight of the whole composition of which the plasticiner represents about 10 to about 30 weight percent based on the weight of the whole composition, and the bloadhesive carrier is in an amount from about 20 to about 44 weight percent based on the weight of the whole composition. More preferably, the composition is comprised of 20 to 34 weight percent of karaya gum, about 20 to 53 weight percent of at least one glycol, and about 10 to 25 weight percent of lideachen base and is further comprised of a binder in or smuleifier an amount sufficient to bind the other impredients.

to a method of administering one or more local anesthetics to a subject in need of such local anesthetic. The term "administering" is intended to mean any mode of application which results in the physical contact of the composition with an anatomical

Another embodiment of the invention relates

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celsius (°C).

site in need of anesthesia. The term "subject" is intended to include all warm-blooded mammals, including humans.

The following examples will further describe
the instant invention, and are used for the purposes
of illustration only, and should not be considered as
limiting in any way the invention being disclosed
herein. Percent (8) as used in these examples refor
to percentage of the liquid formulation on a weight to
10 weight basis and temperatures are given in degrees

Example 1

Ingredient	1 (W/W)
Adhesive (karaya gum) Binder (lecithin) Solvent (propylene glycol) Solvent/plasticizer (glycerin) Amesthetic sgent base (lidocaine base) Amesthetic sgent salt (prilocaine hydrochloride)	21 11 7 19 28 14

The final product is manufactured by first blending the lidecatine bases, prilocatine hydrochlorides, propylace glycol, lecitine and glycerin at about 70 to 90°C until all of the drug is dissolved. The solution is them cooled to 30 to 30°C prior to adding the karge que. Once the karaya que is added, the final composition is applied to a suitable backing material such as a non-woven, polyester film (for example, the film sold under the trademark Sontara 8100, manufactured by DuPort de Nemoure, E.I. and Co., Wilmington, DE) and warmed to about 100°C to accelerate the formation or the gel into its final, finite form.

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29 Example 2

	Ingredient	3_(w/w).
5	Adheeive (karaya gum)	30
	Solvent/plesticizer (glycerin)	30
	Solvent (propylene glycol)	39
	Anesthetic egent base (lidoceine base)	0.7
	Anesthetic agent salt	0.3
10	(prilocaine hydrochloride)	

The procedure set forth in Example 1 is used with appropriate substitutions of quantities to prepare this formulation.

Example 3

	Ingredient	\$_(w/w)
	Adhesive (karaya gum)	21
20	Binder (lecithin)	4
	Solvent (propylene glycol)	3
	Solvent (isocetvl alcohol)	ž
	Solvent/plesticizer (glycerin)	26
	Anesthetic agent bese (lidocaine base)	26
25	Anesthetic egent salt (tetracaine hydrochloride)	13

The procedure of Example 1 is used with appropriate substitution of ingredients to prepare this formulation.

Example 4

	Ingredient	\$_(W/W)
35	Adhesive (karaya gum)	27
	Solvent (propylene glycol)	29
	Solvent/plasticizer (glycerin)	4
	Anesthetic egent base (lidocaine base)	28
	Aneethetio egent salt	12
40	(dyolonine hydrochloride)	

The procedure of Example 1 is used with eppropriete substitution of ingredients to prepare this formulation.

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Example 5

	Ingredient	\$ (w/w)
5	Adhesive (karaya gum) Binder (lecithin) Solvent (propylene glycol)	26 10 7
10	Solvent (butylens glycol) Solvent (butylens glycol) Solvent/plasticizer (glycerin) Anesthetic agent base (lidocaine base) Anesthetic agent salt (dyclonine hydrochloride)	17 10 20 10

The procedure of Example 1 is used with appropriate substitution of ingredients to prepare this formulation.

Example 6

	Ingredient	1 (W/W)
20	Adhesive (karaya gum)	27 12
	Binder (lecithin) Solvent (propylene glycol)	8
	Solvent/plasticizer (glycerin) Anesthetic agent base (lidocaine base)	13 27
25	Anesthetic agent salt (bupivacaine hydrochloride)	13

The procedure of Example 1 is used with appropriate substitution of ingredients to prepare this formulation.

Example 7

	Ingredient	% (W/W)
35	Adhesive (karaya gum)	27
	Binder (lecithin)	12
	Binder (lectural)	8
	Solvent (propylene glycol)	13
40	Solvent/plasticizer (glycerin) Anesthetic agent base (lidocaine base)	13
•••	Anesthetic agent salt (bupivacaine hydrochloride)	27

The procedure of Example 1 is used with 5 appropriate substitution of ingredients to prepare this formulation.

31 Example 8

	Ingredient	\$ (W/W)
5	Adhesive (karaya gum)	21
	Binder (leoithin)	11
	Solvent (propylens glycol)	7
	Solvent/plasticizer (glycerin)	19
	Anesthetic agent base (lidocaine base)	28
10	Anesthetio agent salt (mepivacaine hydrochlorida)	14

The procedure of Example 1 is used with appropriate substitution of ingredients to prepare this formulation.

Example 9

	Ingredient	1 (W/W)
20	Adhesive (Carbopol 934P, a polycarboxylic acid sold by B.F. Goodrich Chemical Company)	20
	Solvent (propylena glycol)	15
	Solvent/plasticizer (glycerin)	20
25	Anesthatic agent base (lidocaine base)	30
	Anesthetic agent salt	15

The procedure of Example 1 is used with 30 appropriate substitution of ingredients to prepare this formulation.

Example 10

35	Ingredient	3_(V/W)
33	Adhasive (karaya gum)	24
	Solvent (propylens glycol)	3
	Solvent/plasticizer (glycerin)	14
	Solvent (isocatyl alcohol)	7
40	Binder (legithin)	4
	Anesthetic agent base (lidocaine base)	32
	Anesthatic agent salt	16

45 The above formulation is prepared by a procedure which is analogous to that set forth in Example 1.

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The addition of up to 2% by weight water in this formulation did not result in precipitation of the anesthetic agent(s) prior to addition of the karaya gus. The addition of 3% to 10% water results in increased precipitation, which at 10% water results in a crystalline mass.

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DAGING A

	Ingredient	4 IMIMI
10	Adhesive (tragacanth gum) Adhesive (pactin)	24 5
	Salvent (propylene glycol)	12 12
15	Solvent/plasticizer (glycerin) Anesthetic agent base (mepivacaine base)	35
15	Anesthetic agent salt (lidocaine hydrochloride)	12

The above formulation is prepared by a procedure analogous to that of Example 1.

Example 12

	Ingredient	3 (8/8/
25	Bioadhesive (karaya gum) Binder (lecithin)	33 9
30	Solvent (propylene glycol) Solvent (dipropylene glycol) Solvent/plasticizer (glycerin) Anesthatic agent base (lidocaine base)	15 17 20

The final product is manufactured by first belanding the lidocaine base, lecithin, propylems glycol, dipropylems glycol and glycerine at about 70 to 90°C until all of the drug is dissolved. The solution is then chilled to about 20 to 40°C prior to adding the karaya qua. Once the karaya qua is added, the final composition is applied to a suitable backing material such as a non-voven polyester film (for example the films sold under the trademark sontate sloo manufactured by DuFout de Nemours, E.I. and Co., Wilmington, DB) and warmed at about 70 to 130°C to scoolerate the formation of the gel into its final

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solid form. This gel can be directly applied to the oral mucosa or overlaid with a skin contact adhesive for skin adhesion.

Example 13

5	Ingredient	\$_(w/w)
	Bioadhesive (karaya gum) Binder (lecithin)	33 5
10	Solvent (propylene glycol) Solvent (dipropylene glycol)	7 12
	Solvent/plasticizer (glycerin)	33
	Anesthetic agent base (lidocaine base)	10

The procedure of Example 12 is used with appropriate substitution of ingredients to prepare this formulation.

Example 14

20	Ingredient	\$ (w/w)
	Bioadhesive (karaya gum)	35
	Binder (legithin)	5
	Solvent (propylene glycol)	7
25	Solvent (dipropylene glycol)	12
25	Solvent/plasticizer (glycerin)	36
	anachatic agent base (lidocaine base)	5

The procedure of Example 12 is used with appropriate substitution of ingredients to prepare this formulation.

Example 15

	Ingredient	1 (W/W)
35	Bioadhesive (karaya gum)	30
	Binder (lecithin)	9
	Solvent (propylene glycol)	6
	Solvent (dipropylene glycol)	15
40	Solvent/plasticizer (glycerin)	15
40	amouthoutic agent hage (lidocaine base)	25

The procedure of Example 12 is used with appropriate substitution of ingredients to prepare this formulation.

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34 Example 16

Ingradient 1 (v/y) hicathesive (karaya gum) 20 hinder (lacithin) 9 solvent (propylene glycol) 10 solvent (propylene glycol) 10 solvent (propylene glycol) 50 Ameribatic agent base (lidocaine base) 40

The procedure of Example 12 is used with appropriate substitution of ingredients to prepare this formulation.

Example 17

	Ingredient	3 (W/W)
20	Bioadhesive (karaya gum)	25 8
	Binder (lecithin) Solvent (isocetyl alcohol)	5
	Solvent (propylene glycol)	12
		10
25	Anesthetic agent base (prilocaine base)	40

The procedure of Example 12 is used with appropriate substitution of ingredients to prepare this formulation.

Example 18

	Ingredient	3 IV/WI
	Bioadhesive (karaya gum)	25
35	Binder (lecithin) Solvent (propylene glycol)	
	solvent (benzvl alcohol)	10
	solvent (dipropylene glycol)	10 5
		40
40	Anesthetic agent base (tetracaine base)	40

The procedure of Example 12 is used with appropriate substitution of ingredients to prepare this formulation.

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35 Example 19

	Ingredient Bioadhesive (karaya gum)	3 (W/W) 30
	Binder (lecithin)	- 8
5	Binder (lecitain)	
	Solvent (propylene glycol)	12
	Solvent (dipropylene glycol)	25
	Solvent (benzyl alcohol)	5
	Solvent (Benzyl alcohol)	
	Solvent/plasticizer (glycerin)	10
10	Anesthetic agent base (dibucaine base)	10

The procedure of Example 12 is used with appropriate substitution of ingredients to prepare this formulation.

Example 20

	Ingredient	\$ (W/W)
	Bioadhesive (karaya gum)	28
20	Bioadhesive (Carbopol 934 Trademark of B.F. Goodrich)	2
	Solvent (propylene glycol)	6
	Solvent (dipropylene glycol)	15
	Solvent/plasticizer (glycerin)	15
25	Binder (legithin)	9
	Anesthetic agent base (lidocaine base)	25

The procedure of Example 12 is used with appropriate substitution of ingredients to prepare this formulation. The only difference is that the carbopol 934 is added to the original blend prior to heating it.

Example 21

35	Ingredient	\$_(w/w)
	Bioadhesive (tragacanth gum)	27
	Bioadhesive (pectin)	6
	Binder (lecithin)	9
40	Solvent (propylene glycol)	6
	Solvent (dipropylene glycol)	15
	Solvent/plasticizer (glycerin)	17
	Anesthetic agent base (lidocaine base)	20

The procedure of Example 12 is used with the solvents and anesthetic agent base added in the initial step followed later by the adhesives addition.

36 Example 22

	Ingredient	
		27
5	Bioadhesive (cellulose acetate)	33
	Solvent (dipropylene glycol) Anesthetic agent base (prilocaine base)	20
	Solvent/plasticizer (glycerin)	10
10	This formulation is prepared	according to
10	the procedure which is analogous to the p	rocedure set
	forth in Example 1.	
	Example 23	
15	Ingredient	1 (w/w)
	Bioadhesive (Xanthan gum)	27
	Bioadhesive (Pectin)	6
	ninder (legithin)	9 6
20	colvent (propylene glycol)	15
20		17
		20
	Anesthetic agent base (1100caine base)	
25	The procedure of Example 12 is f	ollowed with
	the appropriate substitution of ingredier	ts.
	Example 24	
		\$ (w/w)
	Ingredient	
30	Drug (miconazole nitrate)	2
	Solvent (propylene glycol)	67
	Thickener (hydroxymethylcellulose)	1
	Adhesive (karaya gum)	30
35	This formulation is prepared by	y dispersing
	the hydroxymethylcellulose into the propy	lene glycol.
	Once the hydroxymethylcellulose is dispers	ed, the drug
	is added at a temperature between 50 and 8	o'C and mixed
		cooled to
40	until dissolved. The sample is then	the karaya
	approximately 20 to 35°C prior to adding	mulation is
	gum. Once the karaya gum is added, the fo	then the
	applied to a sheet of backing materia	i, then the
	individual dosage forms are cut to the des	rrabie suape

to contain the desired amount of drug.

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	EXHIDIS 23	
	Ingredient	\$ (V/V)
5	Drug (miconazole base) Solvent (dipropylene glycol) Plasticizer (glycerin)	5.0 32.5 32.5
	Adhesive (karaya gum)	30.0
10	Example #25 is prepared just	st as Example #24.
	Example 26	
15	Ingredient	3_(W/W)
	Drug (miconazole base)	5.0 17.5
	Solvent (dipropylene glycol)	30.0
	Plasticizer (glycerin)	7.0
20	Solvent (propylene glycol)	10.5
	Binder (lecithin) Adhesive (karaya gum)	30.0
	Example #26 is prepared jus	st as Example #24.
25	Example 27	
	Ingredient	\$ (W/W)
30	Drug (miconazole base)	10
	Solvent (propylene glycol)	35
	Plasticizer (glycerin)	25
	Adhesive (karaya gum)	30
35	Example #27 is prepared jus	st as Example #24.
	Example 28	
	Ingredient	3_(W/W)
40		
	Drug (clotrimazole)	1.0
	Solvent (propylene glycol)	41.3
	Plasticizer (glycerin)	24.7
	Adhesive (karaya gum)	33.0
45	Example #28 is prepared jus	st as Example #24.
	Example 29	
50	Buccal formulations	containing,
	respectively, 5%, 10%, 20%, and 25%	lidocaine were
	prepared according to the procedu	re of foregoing

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examples. A patch containing no drug (placebo patch) was also used.

The patches were tested on nine human subjects. The patch was applied to the buccal cavity of the mouth and removed after 15 ninutes. The patch was placed on the gingival surface, since the gingival surface was found to be the best site to examine for a dose response relationship.

The extant of anesthesia at 5, 10, 15, 30, 45, and 60 minutes after application was determined by measurement of the extent of anesthesia. The exent of anesthesia was determined by a base line disconfort tolerance limit determined by application of a tip of a periodontal probs, to the treated surface. The patient was saked to determine the depth penetration they could tolerate at the various timed intervals.

Five minutes after initiation of treatment there was no statistical differences in pain toleration between the treatment groups, including the placebo and no-patch.

At ten minutes post application the 281 lidocaine patch produced the greatest mean change in response threshold followed by the 10 and 200 lidocaine patches. There was little difference between the 5% lidocaine and placebo patch. Lidocaine concentrations greater than 5% were necessary to produce a significant increase in pain threshold responses, and there was a distinct trend in dose proportionality in the range of 10% - 25% lidocaine.

The median change in response thresholds for the gingival surface group displayed the same relationship. The 25% lidocaine patch provided the greatest anesthetic effect followed by the 10% and 20% lidocaine patches.

35 When all the sites were combined into one group and the median change from baseline was plotted,

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the graph revealed a dose response profile where the doses appear in order of concentration from 10 to 30 minutes post application. The 29% lidecaine patch provided the greatest increase in response threshold. The 10% and 20% lidecaine patch responses were similar with the 20% lidecaine patch being slightly better.

There were no signs of inflammation, tissue damage, or other adverse effects associated with application of the patches.

Similar studies were conducted in which the patch was applied to the gingival sulcus and the interproximal sulcus.

Certain of the lidocaine preparations were distinguised in that they resulted in the numbness of the teeth, an effect not generally observed with topical anesthetics applied in fluid vehicles.

The foregoing examples are illustrative embodiments of the invention and are merely exemplary. A person skilled in the art may make variations and modification without departing from the spirit and scope of the invention. All such modifications and variations are intended to be included within the scope of the invention as described in this smootification and the arcended claims.

Indeed, the present invention is intended to encompass and be suitable for any pharmaceutically active agent, especially any of the following drugs as the pharmaceutically active agent in the composition:

Analysis anti-inflammatory agents such

as, acetaminophen, aspirin, salicylic acid, methyl salicylate, choins salicylate, dyool salicylate, i-menthol, camphor, mefenamic acid, fluphenamic acid, indomethacin, diclofense, alclofense, lbuprofen, ketoprofen, naprozense, pranoprofen, fenoprofen, sulindac, fenbufen, cildanac, flurbiprofen, indoprofen, protifidic acid, fentiasac, tolmetin,

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like:

tiaprofenic acid, bendazac, bufexamac, piroxicam, phenylbutazone, oxyphenbutazone, clofezone, pentazocine, mepirizole, and the like;

pentazocine, mepirizola, dan terike,
z. Druga having an action on the central
narvous system, for example sedatives, hypnotics,
antiamiety syents, analgesics and amethetics, such
as, chloral, bupremorphine, naloxons, haloperidol,
fluphenarine, pentobarbital, phenobarbital,
secobarbital, amobarbital, cydoharbital, codedne,
lidocalne, tetracalne, dyclomine, dibucaine, cocaine,
procaine, mepivacalne, bupivacalne, stidocaine,
procaine, bensocaine, fentawyl, nicotine, and the

3. Antihistaminics or antialleryic spents when as, diphenhydramine, dimenhydrinate, perphenarine, triprolidine, pyrilamine, chlorvyclimine, bropheniramine, dydrocymine, cyclifine, smolifine, clorprenaline, terfenadine, chlorbeniramine, and the like;

Acetonide anti-inflammatory agents, such as hydrocortisone, cortisone, dexamethasone, fluocinolone, triamcinolone, medrysone, prednisolone, prednisone, halcinonide, flurandrenolide, methylprednisolone, fludrocortisone, corticosterone, paramethasone, betamethasone, ibuprophen, naproxen, fenoprofen, fenbufen, flurbiprofen, indoprofen, suprofen, indomethacin, piroxicam, ketoprofen. diflunisal, methyl salicylic acid, aspirin, salicylate, phenylbutazone, sulindac, nefenanic acid. meclofenamate sodium, tolmetin, and the like;

5. Steroids such as, androgenio steriods, such as, testosterone, methyltestosterone, fluoxymaesterone, estrogens such as, conjugate estrogens, esterified estrogens, estrofipate, 176-estradiol esters such as 176-estradiol

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valerate, equilin, mestranol, estrone, estriol, 17gestradiol derivatives such as 170-athinyl estradiol,
distripustiblestrol, progestational agents, such as,
progesterone, 10-norprogesterone, norethindrone,
norethindrone acetata, malengastrol, chloradinone,
ethisterone, medroxyprogesterone acetate,
hydroxyprogesterone capracte, ethynodiol discetate,
norethynodrel, 17a-hydroxyprogesterone,
dydrogesterone, disabilisterone, ethinylestrenol,
norpestrol, demegaetone, promegastone, megastrol
acetate, and the like:

6. Respiratory agents such as, albutarol, terbutaline, metaproteranol, ritodrina, carbutarol, fenotarol, quinterenol, rimiterol, solmefamel, soteranol, tetroquinol, and the like

- 7. Sympathominetics such as, dopamine, norepinephrine, phanylpropanolamina, phenylephrine, pseudoephedrine, amphetamine, propylhexedrine, arecoline, and the lika;
- 8. Antimicrobial agents including antibacterial agents, antifungal agents, antimycotic agents and antiviral agents; tetracyclines such as, oxytetracycline, penicillins, such as, ampicillin, cephalosporins such as, cefalotin, aminoglycosidas, such as, kanamycin, macrolides such as, erythromycin, chloramphenicol, iodides, nitrofrantoin, anti fungals, such as, clotrimazole, miconazole, chloramphenicol, nystatin, amphotericin, fradiomycin, sulfonamides, sulfacetamida, purrolnitrin, eulfamethazine, sulfadiazine, sulfamerazine, sulfamethizole and sulfieoxazole; antivirals, including idoxuridine; clarithromycin; and other anti-infectives including nitrofurazone, and the like;

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	9.	Antihyperte	nsive	agen	ts suc	h as
clonidine,	α-	methyldopa,	reserp	ine,	syrosin	gopine
rescinnani	ne,	cinnarizine,	hydra	zine,	prazos	in, ar
the like:						

10. Antihypertensive diuretics such as, chlorothiaside, hydrochlorothraside, benedoflumethaside, tripanide, methylclothiaside, penfluxide, hydrothiaside, spironolactone, metolazone, and the like;

11. Cardiotonics such as, digitalis, ubidecarenone, dopamine, and the like;

12. Coronary vanddilators such as, oxymnic nitrates such as, nitroglycerine, isosorbitol dinitrate, erythritol tetranitrata, and pentaerythritol tetranitrata, dipyridamole, dilasep, trapidil, trimetaridine, and the like;

 Vasoconstrictors such as, dihydroergotamine, dihydroergotoxine, and the like;
 β-blockers or antiarrhythmic agents

such as, timolol pindolol, propranolol, and the like;

15. Calcium antagonists and other circulatory organ agents, such as, aptopril,

nitrazepam, meprobamate, phenytoin, and the like;
17. Agents for dizziness such as,
isoprenaline, betahistine, ecopolamine, and the like;

18. Tranquilizers such as, reserprine, chlorpromarine, and antianxiety benzodizepines such as, alprazolam, chlordizespoxide, clorazeptate, halzsepam, oxazepam, prazepam, clorazepam, triazolam, lorazepam, dizzepam, and the like;

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- 19. Antipsychotics such as, phenothiazines including thiopropazate, chiopropazine, triflupromazine, mescridazine, piperracetazine, thioridazine, acetophenazine, piperracetazine, thioridazine, acetophenazine, trifluoperazine, and other major tranquisers such as, chioprethixene, thiothixene, haloperidol, bromperidol, loxapine, and molindone, as well as, those agents used at lower doses in the treatment of namese, vomiting, and the like;
- 20. Muscle relaxants such as, tolperisone, baclofen, dantrolene sodium, cyclobenzaprine;
 - 21. Drugs for Parkinson's disease, spasticity, and acute muscle spasss such as levodopa, carbidopa, anantadine, aposcrphine, bromocriptine, selegiline (deprenyl), trihexyphenidyl hydrochloride, bentropine mesylate, procyclidine hydrochloride, baclofen, diseasam, damtroine, and the like;
 - Respiratory agents such as, codeine, ephedrine, isoproterenol, dextromethorphan, orciprenaline, ipratropium bromide, cromglycic acid, and the like;
 - 23. Non-steroidal hormones or antihormones such as, corticotropin, oxytocin, vasopressin, salivary hormone, thyroid hormone, adrenal hormone, kallikrein, insulin, oxendolone, and the like;
 - 24. Vitamins such as, vitamins A, B, C, D, E and K and derivatives thereof, calciferols, necobalamin, and the like for dermatologically use;
 25. Antitumor agents such as, 5-
 - fluorouracil and derivatives thereof, krestin, picibanil, ancitabine, cytarabine, and the like;
 - Enzymes such as, lysozyme, urokinaze, and the like;
 - Herb medicines or crude extracts such as, glycyrrhiza, aloe, Sikon (<u>Lithospermi radix</u>), and the like;

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28.	Miotics	such	as	pilocarpine,	and	the

like:

29. Cholinergic agonists such as, choline, acetylcholine, methacholine, carbachol, bethanechol,

pilocarpine, muscarine, arecoline, and the like;
30. Antimuscarinic or muscarinic
cholinergic blocking agents such as, atropine,

cholinergio blocking agusta security, as sopplasme, hosatropine, scopplasme, hosatropine, setheropelasme, oyulopantolate, tropicanide, propanthaline, anisotropine, dicyclosline, eucatropine, and the like;

13. Mydratics such as, atropine,

31. Mydriatics such as, attorno, cyclopentolate, homatropine, scopolamine, tropicamide, eucatropine, hydroxyamphetamine, and the like;

32. Psychic energizers such as, 3-(2aminopropy)indole, 3-(2-aminobutyl)indole, and the like;

33. Humoral agents such as, the prostaglandins, natural and synthetic, for example PGE, PGE, and FGF,, and the PGE, analog misoprostol.

34. Antispasmodics such as, atropine, nethantheline, papaverine, cinnamedrine, methscopolanine, and the like;

35. Antidepressant drugs such as, isocarboxarid, phenelrine, tranylopyronine, inipranine, antiriptyline, trisipranine, doxepin, designanine, nortriptyline, protriptyline, amoxapine, saprotiline, trasodone, and the like.

36. Anti-diabetics such as, insulin, and anticancer drugs such as, tamoxifen, methotrexate, and the like;

37. Anorectic drugs such as, dextroamphetamine, methamphetamine, phenylpropanolamine, fenfluramine, diethylpropion, mazindol, phentermine, and the like;

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- 38. Anti-allergenics euch as, antazoline, methapyrilene, chlorpheniramine, pyrilamine, pheniramine, and the like;
- Decongestants such as, phenylephrine, ephedrine, naphazoline, tetrahydrozoline, and the
- like;
 40. Antipyretics such as, aspirin.
 - 40. Antipyretics such as, aepirin, salicylamide, and the like;
 41. Antimigrane agents such ae,
- 41. Antimigrane agents such ae, 10 dihydroergotamine, pisotyline, and the like;
 - 42. Anti-malarials euch as, the 4aminoquinolines, alphaaminoquinolines, chloroquine, pyrimethamine, and the like;
 - 43. Anti-ulcer agente such as, misoprostol, omeprazole, enprostil, allantoin, aldioxa, alcloxa, N-methylscopolamine methylsuflate, and the like;
 - 44. Peptides such as, growth releasing factor, and the like;
- 45. Anti-estrogen or anti-hormone agents 20 such as, tamoxifen or human chorionic gonadotropin, and the like.
 - The drugs mantioned above can be used in combination as required. Moreover, the above drugs may be used either in the free form or, if capable of forming salts, in the form of a salt with a euitable acid or base. If the drugs have a carboxyl group, their estere can be employed.
 - All the drugs used are in solid form at ambient, namely room, temperatures and pressures. However liquid drugs can also be employed to the extent that such drugs, in the forms and amounts used do not undesirably affect the adhesive properties of the carrier.
- The acid mentioned above may be an organic acid, for example, methanesulfonic acid, lactic acid, tartaric acid, fumaric acid, maleic acid, acetic acid,

or ann inorganic acid, for example, hydrochloric acid, hydrokreaic acid, phosphoric acid or sulfuric acid. The base may be an organic base, for example, amonis, tricthylanine, or an inorganic base, for example, sodium hydroxide or potassium hydroxide. The esterm mentioned above may be alkyl esters, anyl esters, aralkyl esters, and the like.

When a drug different than an ansethetic agent is used the solvent selected is one in which the drug is soluble. In generally the polyhydric sloobol can be used as a solvent for a wide variety of drugs, other useful solvents are those known to solubilize the drugs in question.

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CLAIMS

- A flexible, finite, bioadhesive composition for topical application comprising:
- a therapeutically effective amount of at least one pharmaceutically active agent which is in solid form at ambient temperatures and pressures;
 - a pharmaceutically acceptable solvent for the pharmaceutically active agent, in an amount from about 5 to about 70 weight percent based on the weight of the whole composition, said solvent including about 5 to about 50 weight percent of a plasticizer for the bloadheatys:
- in admixture with the pharmaceutically active agent in the solvent, a pharmaceutically acceptable polysaccharide bloadhesive in an amount from about 20 to about 50 weight percent based on the weight of the whole composition;
 - wherein the composition is substantially free of water, substantially water insoluble and selfadhesive, and wherein the pharmaceutically active agent is present in non-oxystallized form in the composition.
 - 2. The composition of claim 1, wherein the pharmaceutically acceptable solvent is in an anount from about 20 to about 33 weight percent based on the weight of the whole composition, of which the plasticizer represents about 10 to about 30 weight percent based on the weight of the whole composition, and the bloadhesive is in an amount from about 20 to about 34 weight percent based on the weight of the whole composition.
 - 3. The composition of claim 1, wherein the pharmaceutically active agent is at least one local anesthetic in an amount of about 10 to about 40 weight percent based on the weight of the total composition.

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- The composition of claim 1, wherein the pharmaceutically active agent is from a class of drugs selected from the group consisting of analysesic antiinflammatory drugs, central nervous system drugs, antihistaminic or antiallergic drugs, acitonide antiandrogenic and estrogenic inflammatory drugs. stercids, respiratory drugs, sympathomimetic drugs, antihypertensive antimicrobial drugs. vasodilators. cardiotonic drugs, coronary vasoconstrictors, beta blocking and antiarrhythemic drugs, calcium antagonistic and other circulatory anticonvulsants, anti-vertigo-tranquilizing drugs, antipsychotic drugs, muscle-reactants drugs, anti-Parkinson drugs, non-steroidal hormones, antihormones, vitamins, anti-tumor, enzymes, medicines or crude extracts, mictics, cholineraic agonists, antimuscarinic or muscarinic cholinergic blocking drugs, mydriatics, psychic energizers. humoral agents, antispasmodic drugs, antidepressants, antidiabetics, anorexic drugs, anti-allergic drugs, decongestants, antipyretics, anti-migraine drugs, antimalarial, antiulcer drugs, peptides, and antiestrogens.
- The composition of claim 4, wherein the antimicrobial drugs is an antifungal agent selected from the group consisting of chlotrinasole, micronarals and chloramphenicol
- 6. The composition of claim 4, in which the pharasceutically active agent is one or more staroids selected from the group consisting of endrogenic staroids, including testestarone; mathyltestostarone; fluoxymasterone; estrogenic staroids, including conjugated estrogens, esterified estrogens, esterified estrogens, estropipate, 178-estradiol, 178-estradiol esters such as 378-estradiol valerate, equilin, mestranol, estrone, estrici) 178- estradiol derivatives such as catcone, estrici) 178- estradiol derivatives such as

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178-ethinyl estradiol; diethylstilbestrol, progestational spents, including progestarone and propositions as under the progestarone spents, proceedings, and as 13-norprogestarone, hydroxyprogesterone separete, 17s-hydroxyprogestarone, contain an onrethindrone, morethindrone acetate, melanguetrol, chlormadinone, ethymodiol diacetate, necessary, dydrogestarone, disemptione, ethinylsstrenol, norgestrol, desegeatione, promegeatione, megestrol, campater and progestarone, and anti-estrogen or anti-androgenic staroids.

7. The composition of claim 3, wherein the ansethetic agent is selected from the group consisting of proceine, lidocaine, prilocaine, mepivacaine, dyclonine, dibucaine, benacoaine, chloroproceine, tetracaine, buylavacaine, and etidocaine and is in the form of the base or an acid-addition salt or both forms.

- The composition of claim 7, wherein the acid-addition salt is hydrochloride.
- The composition of claim 1, wherein the bloadhesive is selected from the group consisting of polyacrylates, polyacrylic acids, guns and celluloses.
- 10. The composition of claim 9, wherein the gum is selected from the group consisting of karaya gum, tragacanth gum, peotin gum, xanthan gum, guar gum, cellulose, and cellulose derivatives.
- The composition of claim 1, wherein the solvent for the anesthetic agent is at least one polyhydric alcohol.
- 12. The composition of claim 11, wherein the polyhydric alcohol is a polyalkylene glycol.
- 13. The composition of claim 12, wherein the glycol is selscted from the group consisting of dipropylsne glycol, propylene glycol, ethylene glycol,

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polyethylene glycol, glycorin, butylene glycol, hexylene glycol, polypropylene glycol, and sorbitol. 14. The composition of claim 1, further comprising a backing material conforming to the size and shape of a single dosage of the composition.

15. The composition of claim 1 comprising about 20 to 34 weight percent of karaya gum, about 20 to 53 weight percent of at least ons glycol, and about 10 to

25 weight percent of lidocaine base and further comprising a binder in an amount sufficient to bind the other ingredients.

16. The composition of claim 15 comprising about 10 weight percent of karaya gum, about 6 weight percent proplate glycol, about 15 weight percent of dipropylane glycol, about 15 weight percent of glycerine, about 25 weight percent of lidocaine base and about 9 weight percent of lectiful.

17. The composition of claim 15, comprising about 31 weight percent of knraya gum, about 7 weight percent of propylene glycol, about 12 weight percent of dipropylene glycol, 33 weight percent of glycerin, about 10 weight percent lidocaine base and about 5 weight percent lecitibin.

18. The composition of claim 1 wherein the parameterical agent comprises a therapeutically effective amount of a first local anesthotic agent in base form and a therapeutically effective amount of a different, local anesthotic agent in acid-addition salt form.

19. The composition of claim 18, wherein the first local nearth-fit agent in base form is selected from the group consisting of procaine, dyclonine, lidocaline, prilocaline, neptwacaine, bennocaine, propoxycaine and chloropyrocaine and the local anaesthatio agent in acid-addition salt form is selected from the group consisting of a dyclonine

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- salt, a prilocaine salt, a tetracaine salt, a bupivacaine salt, a mspivacaine salt, a lidocaine salt, a procaine salt, an etidocaine salt, and a dibucaine salt.
- The composition of claim 21, wherein the acid-addition salt is the hydrochloride.
 - The composition of claim 20, wherein the bloadhesive is selected from the group consisting of polyacrylates, polyacrylic acids, gums and celluloses.
 - 22. The composition of claim 21, wherein the gum is selected from the group consisting of karaya gum, tragacanth gum, pectin gum, xanthan gum and guar gum.

23. The composition of claim 22, wherein the

- solvent for the anesthetic agents is at least one polyhydric alcohol.
- The composition of claim 23, wherein the polyhydric alcohol is a polyalkylene glycol.
 The composition of claim 24, wherein the
- glycol is selected from the group consisting of dipropylene glycol, propylene glycol, ethylene glycol, polyethylene glycol, butylene glycol, hexylene glycol, polypropylene glycol, and sorbitol.
- 26. A method of administering one or more pharmaceutically active agent to a subject comprising the steps of:
 - providing the composition set forth in claim 1; and
 - contacting an area of skin or mucous membrane with the composition to administer the pharmaceutically active agent.
 - 27. The method of claim 26, wherein the pharmaceutically active agent is an anesthetic agent selected from the group consisting of procaine, dyclonine, lidocaine, prilocaine, mepivacaine, benzocaine, propoxycaine, chloroprocaine, tetracaine, bupivacaine, etidocaine, and dibuoaine.

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- 28. The method of claim 27, wherein the anesthetic agent is administered in the form of a free base.
- 29. The method of claim 28, wherein the anesthetic agent is administered in the form of an anid-addition salt.
 - 30. The method of claim 29, wherein the solvent is at least one polyhydric alcohol.
 - 31. The method of claim 30, wherein the polyhydric alcohol is a glycol or cycloalkanepolyol.
 - 32. The method of claim 31, wherein the glycol is selected from the group consisting of dipropylene glycol, propylene glycol, polyethylene glycol, glycerin, butylene glycol, hexylene glycol, polypropylene glycol, ochritol, and sthylene glycol.
 - 31. The method of administering a pharmacoutically active agent of laim 26, wherein the pharmacoutically active spent is a combination of a therapeutically affective amount of a first local anesthatic agent in base form; and a therapeutically effective amount of a different, second local anesthatic agent in an acid-addition salt form.
 - 34. The method of claim 33, wherein the first local anesthetic spent in bese form is selected from the group consisting of proceine, dyclonine, lidocaine, prilocaine, nepivacaine, beancoaine, proposycaine and chloroprocaine and the second local anesthetic agent in acid-addition salt form is selected from the group consisting of a dyclonine salt, a prilocaine salt, a tetracaine salt, a bupivacaine salt, a sepivacaine salt, a lidocaine salt, a procaine salt, a netidocaine salt, and a dibucaine salt.
- 35. The method of claim 34, Wherein the acid-35 addition salt is hydrochloride.

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- 36. The method of claim 35, wherein the bioadhesive is selected from the group consisting of polyacrylates, polyacrylic acids, gums and celluloses.
- The method of claim 36, wherein the gum is selected from the group consisting of karaya gum,
 - tragacanth gum, pectin gum, xanthan gum and guar gum.

 38. The method of claim 37, wherein the solvent
 for the anesthetic agents is at least one polyhydric
 - for the anesthetic agents is at least one polyhydric alcohol.
 - 39. The method of claim 38, wherein the polyhydric alcohol is a polyalkylene glycol or cycloalkanepolyol.
 - 40. The method of claim 39, wherein the glycol or polyol is selected from the group consisting of dipropylene glycol, propylene glycol, ethylene glycol, polyethylene glycol, and sorbitol.
- The composition of claim 1, wherein the pharmaceutically active agent is an anti-microbial agent.
- The composition of claim 41, in which the anti-microbial agent in an antifungal agent.
 - The composition of claim 42 in which the anti-microbial agent is oldtrimazole.
 - 44. The composition of claim 43 in which the anti-microbial agent is miconazole.
 - 45. A composition for topical application comprising a therapeutically effective amount of a first local ansethetic agent in base form and a therapeutically effective amount of a different, second local anesthetic agent in salt form in a flexible, finite, pharmaceutically acceptable adhesive-containing solvent for the first and second local anesthetic agents.
 - 46. The composition of claim 45, wherein the first local anesthetic agent in base form is selected from the group consisting of procaine, lidocaine,

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prilocaine, mepivacaine, dyclonine, dibucaine, benzocaine, propoxycaine, and chloroprocaine.

47. The composition of claim 45, wherein the second local anesthetic agent is selected from the group consisting of a dyclonine salt, a pritocaine salt, a tetracaine salt, a bupivacaine salt, a septivacaine salt, a procaine salt, an etidocaine salt, a procaine salt, an etidocaine salt, a procaine salt, and etidocaine salt, a procaine salt, and etidocaine salt, a procaine salt, and etidocaine salt.

48. The composition of claim 45, wherein the first local anesthetic agent in base form is selected from the group consisting of procaine, dyclonine, lidocaine, prilocaine, mapivacaine, bennocaine, propoxycaine and chicoprocaine and the second local anesthetic agent in salt form is salected from the group consisting of a dyclonine salt, a prilocaine salt, a terracaine salt, a bupivacaine salt, a approcaine salt, a procaine salt, an exidocaine salt, a procaine salt, an exidocaine salt, a procaine salt, an exidocaine salt, a procaine salt, and the salt, and a dibucaine salt, and adulting the salt and the salt the salt a

49. The composition of claim 48, wherein the salt is the hydrochloride.

50. The composition of claim 45, wherein the adhesive is a bloadhesive.

51. The composition of claim 50, wherein the first local anseshetic agent is selected from the group consisting of procaine, lidocaine, prilocaine, mepivacaine, dyclomine, dibucaine, benzocaine, propopyycaine and chloroprocaine.

52. The composition of claim 50, wherein the second local anesthetic agent is selected from the group consisting of a dyolonine salt, a prilocoine salt, a tetracaine salt, a buptwacaine salt, a tetracaine salt, a buptwacaine salt, a mapivacaine salt, a lidocaine salt, a procaine salt, an etdocaine salt, an etdocaine salt, and adibucaine salt.

53. The composition of claim 50, wherein the bicadhesive is karaya gum.

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- 54. A method of delivering local anesthetic agente which comprises the topical administration to a mammal of a composition comprising:
- a therapeutically effective amount of a first local anesthetic agent in base form and
 - a therapeutically effective amount of a different, second local anesthetic agent in salt form in admixture with a flexible, finite.
 - in admixture with a flexible, finite, pharmaceutically acceptable, adhesive; and a solvent in the adhesive for the first and
 - second local anesthetic agents.
 - 55. The method of claim 54, wherein the first local anesthetic spent is selected from the group consisting of processe, dyclonine, lidocaine, prilocaine, mepivacaine, benscoaine, propoxycaine and chioroprocaine and the second local anesthetic agent is selected from the group consisting of a dyclonine salt, a prilocaine salt, a tetracaine selt, a bupivacaine salt, a lidocaine salt, a recease salt, an etidocaine salt, and a dibucaine salt.
 - 56. The method of claim 55, wherein the ealt is a hydrochloride.
 - 57. The method of claim 54, wherein the adhesive is a bloadhesive.
 - 88. The method of claim 57, wherein the first local anesthetic agent is selected from the group consisting of proceine, lidocaine, prilocaine, mapivacaine, dyclonine, dibucaine, bensocaine, propopoyacine and chloroprocaine.
 - 59. The method of claim 57, wherein the second local anesthetic syst is selected from the group consisting of a dyclonine salt, a prilocaine salt, a tetracaine salt, a bupivacaine salt, a mepivacaine salt, a lidocaine salt, a procaine salt, an etidocaine salt, as at lidocaine salt, a procaine salt, an etidocaine salt.

60. The method of claim 57, wherein the bloadheaive is karaya gum.
61. The method of claim 59, wherein the salt is a hydrochloride.

INTERNATIONAL SEARCH REPORT

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		CT MATTER (if several dastification	on symbols apply, indicase all) ⁴	TOO SELECTION	
Int.Cl		A 61 K 9/70 A	al Classification and IPC 1 61 L 15/44		
II. FIELDS	SEARCHED				
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Int.Cl	.5	A 61 K	A 61 L		
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III. DOCUS	CENTS CONSIDERE	D TO BE RELEVANT			
Category*	Citation of D	occupent, ^{EE} with indication, where appo	reprists, of the relevant passages 12	Relevant to Claim No. ¹³	
x	UNIVER	217989 (ERNST MORIT: SITAT GREIFSWALD) 30 document	Z ARNDT January 1985, see the	9	
A	Ep.A. 0250187 (JCHNSON & JOHNSON PRODUCTS INC.) 23 December 1987, see page 3, line 1 - page 4, line 41; pages 7-9, examples 2-4; pages 1,12, examples 6,7				
A	EP,A,0 11 Apr	363224 (BLOCK DRUG 11 1990, see pages 7	CO. INC.) ,8, examples 1,2	1-61	
A	WD,A,8 16 Nov	91074D (INNOVATA BI ember 1989	OMED LTD)	1-61	
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	the document but public golds agency which may then the close to establish also or other special is remaint referring to an or means mount published prior is than the priority da than the priority da the control of the priority da the control of the than the priority da the control of the	community 30 control make of the art which it not mility reference inter preference inter	"I have document published after the interpretation of the property of the property of the published of the control of the published of the published of the published of the control of the published of the control of the published of the publis	cialmed levention he considered to cialmed invention reading step when the recover such docto- tr to a person skilled	
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Mme Daemar FRANK

International Application No. Page 2 PCT/US 92/01730

L DOCUMEN	TS CONSIDERED TO BE RELEVANT Clusters of Document, with Indicate	(CONTINUED FROM THE SECONI ics, where appropriate, of the relevant poss	ofer .	Relevant to Ch	in No.
		PHARMACEUTICAL 68, see the whole docum , lines 17-23; page 18		1-61	
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stional application No. INTERNAT. IAL SEARCH REPORT PCT/US 92/01730 Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet) This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following re- Claims Nos: please see remark because they relate to subject master not required to be searched by this Authority, carmely. Although claims 26-40 and 54-61 are directed to a method of treatment of the human/ animal the search has been carried out and based on the alleged effects of the composition. Claims Nos.: because they relate to parts of the intean extent that no meaningful internation se with the second and third sentences of Rule 6.4/a). Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet) ching Authority found multiple inventions in this international as

est accompanied the payment of additional search fees.

Remark on Protest

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

US 9201730 58216

SA

Patent document cited in search report	Peblication date	Pate	st family mber(s)	Publicatio date
DD-A- 217989		None		
EP-A- 0250187	23-12-87	US-A- AU-A- JP-A- US-E-	4713243 7415587 63019152 RE33093	15-12-87 17-12-87 26-01-88 17-10-89
EP-A- 0363224	11-04-90	AU-A- CA-A- JP-A-	4265689 2000277 2196717	12-04-90 07-04-90 03-08-90
WO-A- 8910740	16-11-89	None		
LU-A- 52460	25-06-68	BE-A- DE-A- FR-M- G8-A-	690383 1617282 6733 1108837	29-05-67 06-02-75 24-02-69
		NL-A-	6616878	31-05-67